of the facts that warrant the return of the substance to the United States along with an authorization from the country of export will be included with the request. DEA will evaluate the request after considering all the facts as well as the exporter's registration status with DEA. The substance may be returned to the United States only after affirmative authorization is issued in writing by DEA.

(c) Notwithstanding the time limitations included in paragraph (a) of this section, a registrant may obtain a special waiver of these time limitations in emergency or unusual instances; provided that a specific confirmation is received from the Administrator or his delegate advising the registrant to proceed pursuant to the special waiver.

[36 FR 7815, Apr. 24, 1971, as amended at 37 FR 15923, Aug. 8, 1972. Redesignated at 38 FR 26609, Sept. 24, 1973, and amended at 45 FR 74715, Nov. 12, 1980; 51 FR 5319, Feb. 13, 1986; 52 FR 17290, May 7, 1987; 62 FR 13969, Mar. 24, 1997]

§ 1312.28 Distribution of special controlled substances invoice.

The required five copies of the special controlled substances export invoice, DEA (or BND) Form 236, will be distributed as follows:

- (a) Copy 1 shall accompany the shipment and remain with the shipment to its destination.
- (b) Copy 2 shall accompany the shipment and will be detached and retained by appropriate customs officials at the foreign country of destination.
- (c) Copy 3 shall accompany the shipment and will be detached by the District Director of the U.S. Customs Service at the port of exportation, who shall sign and date the certification of customs on such Copy 3, noting any changes from the entries made by the exporter, and shall then promptly forward Copy 3 to the Drug Control Section of the Administration.
- (d) Copy 4 shall be forwarded, within the time limit required in \$1312.27 of this part, directly to the Drug Enforcement Administration, Drug Operations Section, Washington, DC 20537. The documentation required by \$1312.27(b)(4) of this part must be attached to this copy.

(e) Copy 5 shall be retained by the exporter on file as his record of authority for the exportation.

[36 FR 7815, Apr. 24, 1971, as amended at 36 FR 13387, July 21, 1971. Redesignated at 38 FR 26609, Sept. 24, 1973, and amended at 45 FR 74715, Nov. 12, 1980; 51 FR 5319, Feb. 13, 1986; 52 FR 17291, May 7, 1987; 53 FR 48244, Nov. 30, 1988; 62 FR 13969, Mar. 24, 1997]

§1312.29 Domestic release prohibited.

An exporter or a forwarding agent acting for an exporter must either deliver the controlled substances to the port or border, or deliver the controlled substances to a bonded carrier approved by the consignor for delivery to the port or border, and may not, under any other circumstances, release a shipment of controlled substances to anyone, including the foreign consignee or his agent, within the United States.

§ 1312.30 Schedule III, IV, and V nonnarcotic controlled substances requiring an import and export per-

The following Schedule III, IV, and V non-narcotic controlled substances have been specifically designated by the Administrator of the Drug Enforcement Administration as requiring import and export permits pursuant to sections 1002(b)(2) and 1003(e)(3) of the Act (21 U.S.C. 952(b)(2) and 953(e)(3)):

- (a) Dronabinol (synthetic) in sesame oil and encapsulated in a soft gelatin capsule in a U.S. Food and Drug Administration approved product.
 - (b) [Reserved]

 $[52~{\rm FR}~17291,~{\rm May}~7,~1987,~{\rm as}~{\rm amended}~{\rm at}~64~{\rm FR}~35930,~{\rm July}~2,~1999]$

TRANSSHIPMENT AND IN-TRANSIT
SHIPMENT OF CONTROLLED SUBSTANCES

§ 1312.31 Schedule I: Application for prior written approval.

- (a) A controlled substance listed in schedule I may be imported into the United States for transshipment, or may be transferred or transshipped within the United States for immediate exportation, provided that:
- (1) The controlled substance is necessary for scientific, medical, or other legitimate purposes in the country of destination, and

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- (2) A transshipment permit has been issued by the Administrator.
- (b) An application for a transshipment permit must be submitted to the Drug Enforcement Administration, Drug Operations Section, Washington, DC 20537, at least 30 days, or in the case of an emergency as soon as practicable, prior to the expected date of importation, transfer or transshipment. Each application shall contain the following:
 - (1) The date of execution;
- (2) The identification and description of the controlled substance;
 - (3) The net quantity thereof;
- (4) The number and size of the controlled substance containers;
- (5) The name, address, and business of the foreign exporter;
- (6) The foreign port of exportation;
- (7) The approximate date of exportation:
- (8) The identification of the exporting carrier;
- (9) The name, address and business of the importer, transferor, or transshipper;
- (10) The registration number, if any, of the importer, transferor or transshipper;
 - (11) The U.S. port of entry;
 - (12) The approximate date of entry;
- (13) The name, address and business of the consignee at the foreign port of entry:
- (14) The shipping route from the U.S. port of exportation to the foreign port of entry;
- (15) The approximate date of receipt by the consignee at the foreign port of entry; and
- (16) The signature of the importer, transferor or transshipper, or his agent accompanied by the agent's title.
- (c) An application shall be accompanied by an export license, permit, or a certified copy of the export license, permit, or other authorization, issued by a competent authority of the country of origin (or other documentary evidence deemed adequate by the Administrator).
- (d) An application shall be accompanied by an import license or permit or a certified copy of such license or permit issued by a competent authority of the country of destination (or

- other documentary evidence deemed adequate by the Administrator), indicating that the controlled substance:
- (1) Is to be applied exclusively to scientific, medical or other legitimate uses within the country of destination;
- (2) Will not be exported from such country; and
- (3) Is needed therein because there is an actual shortage thereof and a demand therefor for scientific, medical or other legitimate uses within such country.
- (e) Verification by an American consular officer of the signatures on a foreign import license or permit shall be required, if such license or permit does not bear the seal of the authority signing them.
- (f) The Administrator may require an applicant to submit such documents or written statements of fact relevant to the application as he deems necessary to determine whether the application should be granted. The failure of the applicant to provide such documents or statements within a reasonable time after being requested to do so shall be deemed to be a waiver by the applicant of an opportunity to present such documents or facts for consideration by the Administrator in granting or denying the application.
- (g) The Administrator shall, within 21 days from the date of receipt of the application, either grant or deny the application. The applicant shall be accorded an opportunity to amend the application, with the Administrator either granting or denying the amended application within 7 days of its receipt. If the Administrator does not grant or deny the application within 21 days of its receipt, or in the case of an amended application, within 7 days of its receipt, the application shall be deemed approved and the applicant may proceed.

[36 FR 7815, Apr. 24, 1971, as amended at 37 FR 15923, Aug. 8, 1972. Redesignated at 38 FR 26609, Sept. 24, 1973, and further amended at 45 FR 74715, Nov. 12, 1980; 51 FR 5319, Feb. 13, 1986; 53 FR 48244, Nov. 30, 1988; 62 FR 13969, Mar. 24, 1997]